

CACHAÇA DISTILLATION INVESTIGATED ON THE BASIS OF MODEL SYSTEMS

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Abstract - This work reports experimental tests using commercial spirits that were diluted and had their initial composition modified in order to better measure the distillation behavior of selected minor compounds of importance for the quality of alcoholic beverages. Such compounds were added to the commercial *cachaça* and the corresponding model wine, obtained after the spirits' dilution, was distilled. In this way a more precise distillation profile could be determined for those minor compounds. The alembic heating was performed by electrical resistance and the corresponding heat transfer coefficient was determined by analyzing the thermal behavior of the still during the distillations. A simulation algorithm was developed, including the mass and enthalpy balances as well as the heat transfer flux to the boiling wine. Good agreement between experimental and simulated results was obtained for the distillate rate, alcoholic graduation, temperature and most minor component profiles.

Keywords: *Cachaça*; Distillation; Simulation; Heat transfer coefficient.

INTRODUCTION

Brazilian sugar cane spirit, also called "*cachaça*", is a beverage obtained from the distillation of fermented sugar cane juice (wine), with an alcoholic graduation between 38% v/v and 48% v/v (Brasil, 2005). It is the world's third most consumed spirit by volume, and its consumption is increasing on the international market because of its exotic and special flavor. Nowadays, Brazilian production of sugar cane spirit is estimated at 1.3 billion liters per year.

Cachaça is produced by fermentation of the sugar cane juice, followed by distillation and aging. Fermentation exerts the main influence on product quality, because minor components, such as alcohols, organic acids, carbonyl compounds and

esters, are generated during this step and are responsible for the spirits flavor (Lurton *et al.*, 1995).

Ethanol is responsible for the beverage's body. Higher alcohols, such as isoamyl alcohol, represent the main group in terms of the spirits' flavor (Nykänen, 1986). Some compounds can be considered to be indicators of *cachaça* quality. The propanol level is usually low in spirits of good quality (Almeida and Barreto, 1972). The acetaldehyde concentration must be low, as it is related to the "hangover" syndrome (Nascimento *et al.*, 1997). The methanol level should be controlled because it causes severe intoxication.

Cachaça quality is also influenced by the distillation, since in this step ethanol is concentrated

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in the distillate and the level of minor compounds is simultaneously determined. The traditional distillation of wine is carried out in a pot still working as a differential distillation unit. The main cut of the distillate, the heart, is concentrated in ethanol and corresponds to the final product (Yokoya, 1995). Two further cuts are usually obtained, the head, which contains the largest amounts of the minor compounds with higher volatility (acetaldehyde, methanol and ethyl acetate), and the tail, used for recovering the residual ethanol still present in the wine.

Although ethanol distillation is a widely investigated subject, the influence of minor compounds on the distillation path is an almost ignored theme. In a previous work of our research group (Scanavini *et al.*, 2009), *cachaça* distillation was investigated by conducting experiments in a small scale alembic heated by direct fire and taking into account seven minor components. The experimental runs were reproduced by simulation based on a differential distillation algorithm and a good agreement was achieved between the experimental and the simulation results in terms of temperature and alcoholic graduation profiles, and also heart concentrations of the main congeners.

Other works reported the use of computer simulation to study beverage distillation. Osório *et al.* (2004, 2005) worked with *Pisco*, a distilled drink typical of Chile and Peru, optimizing the process in terms of the preferences of enologists and the best operational conditions of the distillation column. They included in the simulations six to eight minor components, but only in the latter work did they carry out a comparison of the simulated results with a very limited set of experimental data. In fact, they compared only the recoveries of three components in the heart fraction, ethanol and two minor ones. Gaiser *et al.* (2002) discussed the use of the commercial software Aspen Plus for simulating the production of grain whisky, typically manufactured in a continuous distillation. Batista and Meirelles (2011) simulated an industrial unit for continuous *cachaça* distillation, taking into account ten minor components and performed a comparison with results of an industrial equipment. Zavala-Loria *et al.* (2011) worked with an alcoholic mixture and proposed a thermodynamic model, considering reflux and efficiency, to simulate batch distillation of ideal and non-ideal mixtures.

In the present work a new set of experimental tests is reported using commercial spirits that were diluted and modified by the addition of minor components in order to better measure their

distillation behavior in a model wine. In this way, additional effects that could occur in the distillation of a real fermented must were excluded, as for instance methanol generation associated with the presence of methoxylated pectin. On the other hand, the selected model system is sufficiently complex to represent a real wine in an approximate way, since it contains all minor components important for the quality of spirits. The alembic heating was performed by electrical resistance and the corresponding heat transfer coefficient was determined, allowing the calculation of the vaporization rate in the simulation algorithm. The experimental and simulated results were compared for the alcoholic graduation, temperature and for all the minor component profiles. The comparison was also performed for the head and heart cuts. Such a detailed comparison of experimental and simulated data helps to highlight the aspects to be improved in the use of simulation tools for studying spirit distillation and its quality. In fact, the reliable use of simulation tools for describing purification processes applied to products whose quality is related to minor components of utmost importance still requires further investigation. This is the case of alcoholic beverages, but also of aromas, cosmetics, perfumes, and edible oils.

MATERIAL AND METHODS

Experimental Trials

In order to study the thermal behavior of the still, five experiments were performed using different solutions: (1) distilled water; (2) 96% v/v ethanol and (3) three diluted sugar cane spirits (*cachaça*), with approximately 9% v/v ethanol. The still was thermally insulated, using a thermal blanket, in order to avoid heat losses to the environment. A scheme of the insulated still is shown in Figure 1. It was considered that the insulation does not allow any significant heat exchange between the equipment and the surroundings.

The choice of solutions for the two first experiments was made by considering that both water and ethanol (96% v/v) would not change composition during the distillation, that is, any sample of distillate obtained would have the same composition. For this reason it was not necessary to analyze the composition along the distillation path and the heat transfer coefficient could be easily evaluated.

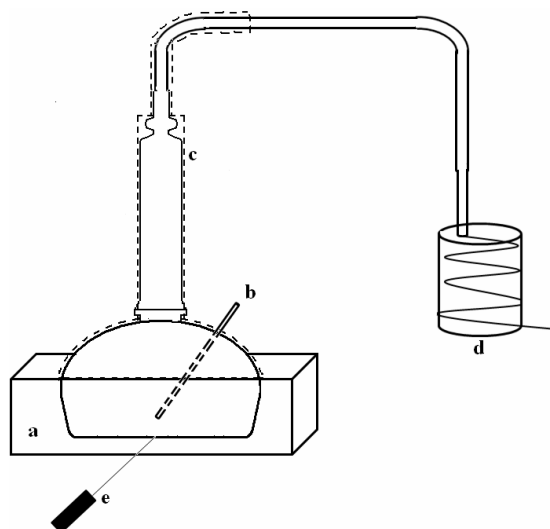


Figure 1: Simple still: (a) heat system; (b) thermometer; (c) the dashed line represents the thermal blanket used for insulation; (d) condenser (e) temperature sensor under the still.

In order to improve the description of the minor compounds distillation, and also to analyze the thermal behavior of a real distillation, experiments were performed with an already distilled spirit, which was diluted and had its initial composition modified. For these experiments, multidistilled commercial *cachaça* was used.

Firstly, those *cachaça* samples were analyzed in order to determine the alcohol content, volatile acidity and composition. On the basis of these values, the composition modification and dilution were conducted. The modification of composition was done through the addition of minor compounds (acetaldehyde, acetic acid, ethyl acetate, isoamyl alcohol and methanol) in known concentrations. These concentrations were defined based on the range of values for wine compositions found in the literature (Oliveira, 2001; Oliveira *et al.*, 2005; Scanavini *et al.*, 2009) and the limits of concentrations established by the Brazilian legislation for *cachaça* (Brasil, 2005).

The values are presented in Table 1.

The aim of modifying the spirits' composition was to obtain concentrations of minor compounds of an order of magnitude close to the maximum values reported in the literature (Table 1), so that their distillation paths could be followed with higher precision.

Three experiments with three different samples of the same brand of *cachaça* were performed. In case of the first wine (W_1), the *cachaça* sample was diluted and modified in its acetaldehyde, ethyl acetate, methanol and acetic acid contents. As can be seen in Table 1, the contents of almost all minor compounds were close to the maximum values reported in the literature. The only exception is ethyl acetate, where the composition modification exceeded the maximum value. In the case of wines W_2 and W_3 , the *cachaça* samples were diluted and modified in their contents of methanol, acetic acid and isoamyl alcohol.

Table 1: Brazilian Legislation for *cachaça* and wine compositions

Compound	Cachaça Legislation (mg/100 mL AA) ^a	Wine Composition in the literature (mg/100 mL AA) ^b	W_1^c	W_2^c	W_3^c
Alcohol Content (% v/v)	38 – 48	8 – 10	8.88	8.80	8.72
Volatile acidity (in Acetic Acid)	≤ 150.0	7.1 – 1152.0	1062.68	1209.99	1162.47
Acetaldehyde	≤ 30.0	6.6 – 96.1	97.79	2.69	9.33
Ethyl Acetate	≤ 200.0	6.0 – 14.4	638.32	123.25	52.94
Methanol	≤ 20.0	ND	32.17	21.57	34.56
n-propanol	NS ^d	7.2 – 78.4	82.96	50.35	73.34
Isobutanol	NS	16.0 – 56.9	92.63	53.89	85.66
Isoamyl Alcohol	NS	31.4 – 218.0	153.25	232.67	219.50
Sum of Higher Alcohols	≤ 360.0	31.4 – 329.0	328.84	336.91	378.50

^a AA = Anhydrous Alcohol; ^bOliveira, 2001; Oliveira *et al.*, 2005; Scanavini *et al.*, 2009;

^c Cachaças after composition modification and dilution, compositions also in (mg/100 mL AA); ^d NS = not specified.

It should be observed that the initial acetic acid and methanol contents were modified in all wines. The initial concentrations of the minor components with higher volatility (acetaldehyde and ethyl acetate) were modified only in W_1 . The initial concentration of the main higher alcohol (isoamyl alcohol) was modified in W_2 and W_3 .

During the experiments, flow rates were measured over the time necessary to obtain 50 mL of distillate. These flow rates, in mL/s, were transformed into molar flows (mol/s) and subsequently into rates of heat transfer Q (J/s), on the basis of the enthalpies of vaporization (in J/mol) evaluated at the boiling temperatures of each collected sample.

In the case of the diluted spirits, the distillation of each compound important to the spirit quality was monitored by GC analysis, density and volatile acidity. These analyses were conducted for each 100 mL sample obtained by combining two subsequent distillate samples of 50 mL.

The following temperatures were measured during the experiments: the temperature inside the still (the boiling temperature of the solution, T_b), the room temperature, the resistance temperature (T_R) and also the temperature under the still during distillation (T_u). This last temperature represents the temperature of the exterior side of the metal sheet at the still bottom. The resistance temperature was kept constant at 318°C in all experiments.

Physical and Chemical Analysis

The alcohol content was determined by measuring the mixture relative density at 20°C, in a Anton Paar densimeter. Considering that the minor compounds represent a maximum of 0.7% in mass of the collected samples, they do not interfere significantly in the density values and the corresponding alcoholic graduations changed no more than 0.2 °GL as a consequence of their presence, so that measuring the alcohol content through density is a reliable procedure. The volatile acidity was determined by the volumetric method suggested by Amerine and Ough (1974). According to this method, the wine or distillate sample must be further distilled under direct steam injection so that the volatile acidity is withdrawn from the sample and then titrated using a standard NaOH solution. The volatile acidity was calculated and expressed in terms of acetic acid. The other minor compound compositions were determined by gas chromatography (Shimadzu GC 17 A gas chromatograph), with flame ionization detector

(FID), on a DB-WAX column (30m x 0,25 mm x 0,25 mm), with an AOC-20i automatic injector. The method was based on the one described in Oliveira *et al.* (2005): Helium was used as the carrier gas, the linear velocity kept constant at 30 cm/s with a temperature ramp of 35°C for 8 min, then an increase to 65°C at a rate of 1°C/min and an increase to 200°C at a rate of 30°C/min, remaining at the final temperature for 5 min; the split ratio was 1:50 and injection volume 1.5 µL. External standard curves were used to determine the compound amounts.

Simulation Program

The simulation program was based on the algorithm suggested by Scanavini *et al.* (2005) and by Ceriani and Meirelles (2004). The batch distillation in the alembic was considered to be a multicomponent differential distillation without the occurrence of reflux. In the case of the vapor-liquid equilibrium calculations, it was assumed that the liquid and vapor phases are nonideal. The activity coefficients were calculated by the NRTL equation and the fugacity coefficients were evaluated using the virial equation and the Hayden O'Connell approach, as described elsewhere (Scanavini *et al.*, 2009).

The algorithm used is based on the following set of Equation:

Enthalpy balance in the still bottom:

$$Q = (UA) \cdot (T_u - T_b) \quad (1)$$

$$Q = v \cdot H_v + L \frac{dH_L}{dt} \quad (2)$$

Mass balance in the still bottom:

$$\frac{dL}{dt} = -v \quad (3)$$

$$\frac{d(Lx_i)}{dt} = -v \cdot y_i \quad (4)$$

Equilibrium relationship:

$$\phi_i \cdot y_i \cdot P = \gamma_i \cdot x_i \cdot P_{vpi} \quad (5)$$

Mass balances in the distillate receiver:

$$\frac{dD}{dt} = v \quad (6)$$

$$\frac{dD_i}{dt} = v \cdot y_i \quad (7)$$

where Q is the heat transferred to the boiling wine (J/s), T_u is the heat source temperature evaluated from the temperature under the still ($^{\circ}\text{C}$), T_b is the wine boiling temperature ($^{\circ}\text{C}$), U is the heat transfer coefficient ($\text{W}/\text{m}^2\text{C}$), A is the heat transfer area (m^2), v is the vaporization rate (mol/s), H_v is the vapor phase enthalpy (J/mol), L is the total amount of liquid in the still (mol), H_L is the liquid phase enthalpy (J/mol), t is time (s), x_i is the liquid phase mole fraction of component i , y_i is its vapor phase mole fraction, ϕ_i and γ_i are the fugacity and activity coefficients, respectively, P is the total pressure (Pa), P_{vpi} is the vapor pressure of component i (Pa), D is the total amount of distillate (mol) and D_i is the total amount of component i in the distillate (mol).

RESULTS AND DISCUSSION

Figure 2 shows the curves of the wine temperature from the beginning of heating, the vaporization rate, in mol/s, and the values of UA for experiment W₂. During the distillate withdrawal, the boiling temperature does not show a large variation, but increases continuously, as a consequence of the decreasing concentration of ethanol in the wine. The vaporization rate remains almost constant along the distillation period. The behavior observed for the UA-values shows a slightly larger variation, but the measured changes around the average value were never higher than $\pm 5\%$.

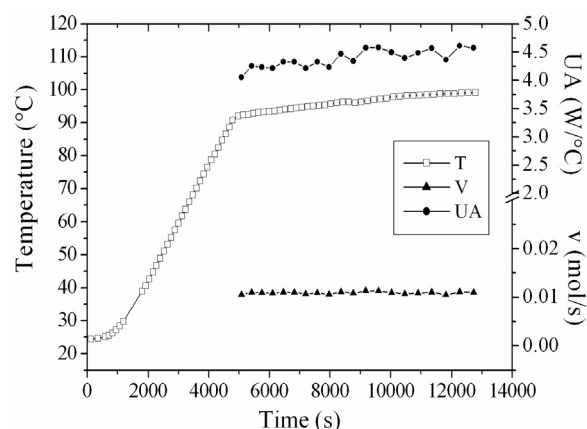


Figure 2: Profiles of temperature, vaporization rate (v) and UA-values for experiment W₂.

The UA-values were estimated on the basis of the measured vaporization rates for all experimental

runs. In the case of the diluted spirits distillations, the effect of the wine and distillate changing compositions were taken into account through the wine boiling temperatures and the distillate heat of vaporization.

The calculated UA-values depend on the specified temperature difference between the heat source and the wine temperature. Industrial alembics are normally heated by direct fire or steam. In small scale experiments, the use of electrical heating makes it easier to control the heat transfer and, consequently, the vaporization rate. In each one of these cases, the heat source temperature will be different, affecting the estimated UA-values (see Eq. (1)). In order to avoid this problem, the heat source temperature was assumed to be the temperature of the exterior side of the still bottom (T_u), as measured in the experimental runs. In the case of the water boiling, this temperature was 202°C and for the hydrated alcohol or diluted spirits distillations, it was equal to 197°C . The calculated UA-values are given in Table 2.

Table 2: Values of the heat transfer coefficients UA

Experimental run	UA (W / $^{\circ}\text{C}$)
Water	4.84 ± 0.23
Ethanol	4.07 ± 0.15
W ₁	4.35 ± 0.20
W ₂	4.32 ± 0.16
W ₃	4.31 ± 0.14

Thus, simulations were carried out in order to reproduce the experiments with wines W₁, W₂ and W₃. In these simulations, an average value for UA calculated on the basis of the results shown in Table 2 was used ($4.38 \text{ W}/^{\circ}\text{C}$).

Figure 3 gives the alcoholic graduation profiles and indicates that the simulation provided results very close to the experimental alcohol content. The deviations between experimental and simulated values are shown in Table 3.

Figure 3 also shows the temperature profile for the distillation of wine W₃. Table 4 gives the average deviations and also the initial (T_{initial}) and final (T_{final}) temperatures obtained in each case. The simulated values are very close to the experimental ones, presenting a maximum deviation of 0.70% (W₃). The initial boiling temperatures of the mixtures gave the highest deviations among all points, with values between 2.0 and 2.7% .

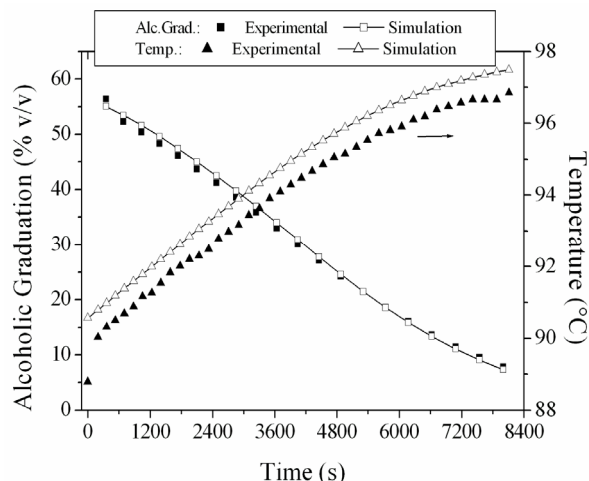


Figure 3: Distillate alcoholic graduation and wine temperature profiles for W_2 and W_4 distillations, respectively.

Table 3: Distillate alcoholic graduation in the experimental and simulated runs

	Experiment	GL _{initial}	GL _{final}	Average Deviation (%) ^a
W_1	Experimental	56.6	7.6	7.2
	Simulation	54.0	8.7	
W_2	Experimental	56.4	7.8	5.4
	Simulation	54.9	8.6	
W_3	Experimental	56.1	7.3	4.6
	Simulation	54.7	7.9	

^aAverage Deviation: $AD = \frac{100}{N} * \sum_{n=1}^N \frac{|r_{exp} - r_{sim}|}{r_{exp}}$, where N is the total number of experimental points, r_{exp} is the experimental result and r_{sim} the simulated one.

Table 4: Boiling temperatures in the experimental and simulated runs

	Experiment	T _{initial} (°C)	T _{final} (°C)	Average Deviation (%)
W_1	Experimental	88.5	96.9	0.64
	Simulation	90.3	97.3	
W_2	Experimental	88.8	96.7	0.68
	Simulation	90.5	97.3	
W_3	Experimental	88.8	96.9	0.70
	Simulation	90.6	97.4	

The mass concentration profiles of each component considered in the experimental runs and in the simulations are shown in Figures 4 to 7. In general, a good proximity was observed between the experimental and simulated concentration profiles, with the exception of isoamyl alcohol and acetic acid.

The most volatile compounds, acetaldehyde and ethyl acetate, shown in Figure 4, had distillation profiles that were well described by the simulations, but the initial experimental values were always lower than the simulated ones. In the case of acetaldehyde, it is possible that there was some loss of this compound by evaporation during the condensation of

the distillate fractions. The boiling point of acetaldehyde is very low ($T_b = 21^\circ\text{C}$), and the condensation was carried out using only running water for cooling.

Figure 5 shows the results for the methanol profiles. For this compound, the simulation results indicate higher distillate concentrations in comparison with the experimental values at the beginning of the distillation, which decreases along the process, with values changing from 8×10^{-5} to 3×10^{-5} g/g. In the experiments, the concentrations also decrease through the process, but starts at a higher value, of the order of 10^{-4} g/g.

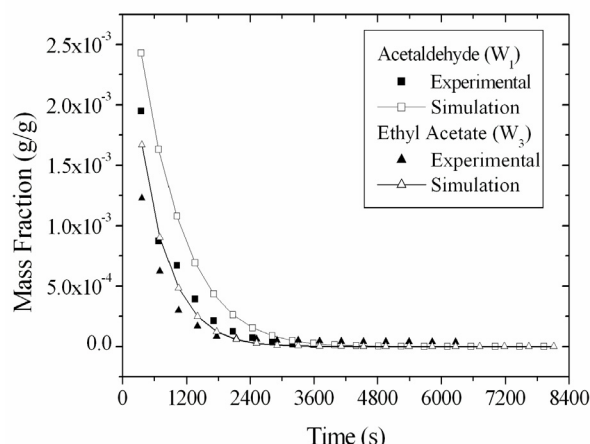


Figure 4: Distillate acetaldehyde and ethyl acetate profiles during distillation.

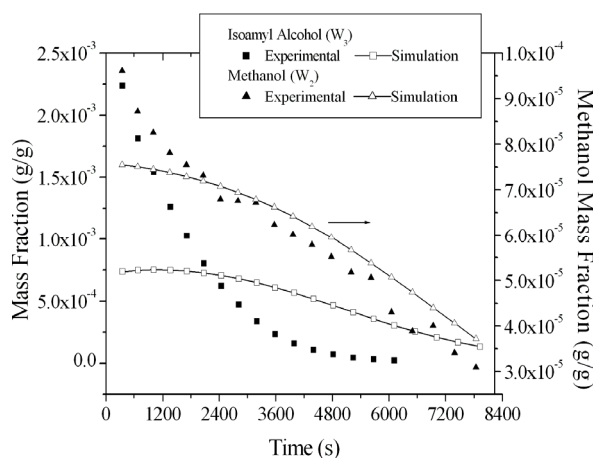


Figure 5: Distillate methanol and isoamyl alcohol profiles during distillations.

According to literature reports (Madrera and Alonso, 1996; Cortés *et al.*, 2002; Cortés *et al.*, 2005; Claus and Berglund, 2005), methanol distillate concentrations showed an unexpected increase in concentration at the end of the distillation. This behavior is different than would be expected due to methanol's high volatility and low boiling point: it should distill mainly at the beginning and its concentration steadily decreases along the distillation path. Despite the somewhat larger differences observed in our results between methanol's experimental and simulated concentration values, in comparison to the prior profiles, the results for methanol are still in good agreement and do not confirm those literature reports of an unexpected increase in its concentration at the end of the distillation. Considering that those literature works

investigated alembic distillation of real fermented musts, a possible explanation of the observed behavior might be an eventual influence of other compounds on methanol volatility or an eventual methanol generation by, for instance, pectin degradation during real wine distillation.

These three compounds, as the most volatile ones, are mostly withdrawn in the head fraction. As said previously, it is important to separate methanol and acetaldehyde from the spirits since they are undesirable for *cachaça* quality. Normally, the head cut used for extracting these compounds from the product corresponds to 1 - 2% of the volume added to the still bottom. This means that, in these experiments, a volume between 72 and 144 mL should be separated. But the analyses of the samples were done every 100 mL, so the head cut was

contained in for the first 200 mL of distillate. For the experimental results, if a volume in this range is separated, this would be enough to withdraw the largest amount of these compounds in the head fraction.

For ethyl acetate in W_1 , the experiment in which this compound's initial concentration was modified, 200 mL of head was not enough to make the spirit conform the Legislation, since a content of 205.8 mg/100 mL AA was obtained.

Considering that, when 200 mL was taken as the head fraction, this was not enough to reduce the contents of some lighter components in the heart fraction to within the required values, such as in W_1 (ethyl acetate), the simulation results can be used as a tool to determine when the cut should be done in order to distill the spirit according to the Legislation limits.

The higher alcohols propanol and isobutanol did not have their initial contents modified as they were already close to the maximum values found in the wine compositions reported in the literature. Their profiles are shown in Figure 6. For isobutanol, the simulation showed higher values right from the beginning, except for the first experimental point; but the experimental and simulation results are very close. It should be noted that isobutanol concentrations at the end of the distillation have values below the experimental detection limit. The n-propanol profile was even better described by the simulation, although in the beginning the simulation results are higher and in the end the curves invert the prior behavior.

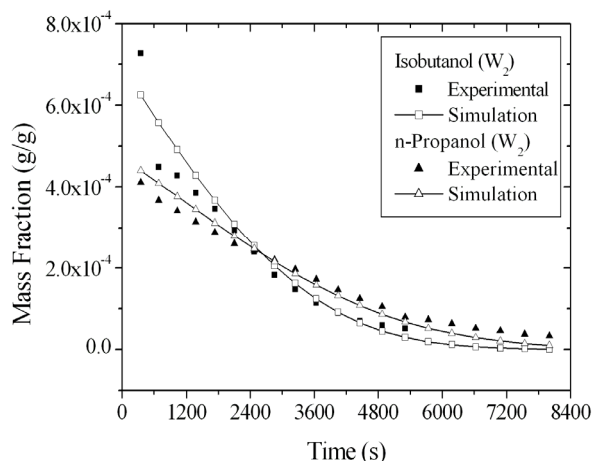


Figure 6: Distillate isobutanol and n-propanol profiles for W_2 distillation.

As pointed out before (Table 1), there are no specific Legislation limits for these two higher

alcohols in separate. So the cuts should be analyzed for all the higher alcohols together, which also includes isoamyl alcohol.

Figure 5 also shows the distillate profiles of isoamyl alcohol. The experimental curve of this higher alcohol crosses the simulation results and presents lower final concentrations. The larger differences between simulation and experiments could be attributed to an inappropriate prediction of isoamyl alcohol volatility. According to the experimental profiles, it can be seen that this higher alcohol is actually more volatile than the simulation predicts. In fact, despite its relatively large carbon chain, isoamyl alcohol is a light component in a water-rich environment, such as alcoholic wine. In such an environment, this compound's activity coefficient is very large and probably its value was not well predicted by the NRTL equation. According to the simulation results, isoamyl alcohol should be more accumulated in the heart fraction than indicated by the experimental results. Different experimental data sets can be found in the literature for the binary mixture isoamyl alcohol and water (Gmehling and Onken, 1981; Cho *et al.*, 1984; Dechema online, 2010) and, from a quantitative point of view, they do not always agree. In fact, those data exhibit relatively large differences in the region of low isoamyl alcohol concentration, the region of interest in the present case. The set of parameters available in the Aspen Plus database was used in the present work and considered to be the most reliable source at the time. Although alternative sources of parameters could be tested and eventually allow lower deviations between experimental and simulated distillation results, such a decrease in the deviations does not discriminate between errors due to the phase equilibrium data and those related to the distillation process itself. In fact, compensation of errors could occur in such a case, suggesting that new equilibrium data for this binary mixture are probably required to solve the conflict between the already available experimental data.

For the volatile acidity, measured in terms of acetic acid (Figure 7), the experimental and simulation curves showed the largest deviations. The experimental data are of the order of 10^{-4} , while the simulations are of an order of 10^{-5} . Despite the differences between the curves, it should be noted that the simulation results show a trend of increasing acetic acid concentration, the actual behavior observed during the experimental distillation. Among the minor compounds considered, this acid is the least volatile, so its content increases more at the end of the distillation, being more concentrated in

the tail fraction. For this minor compound, the heart cut is the one that determines its quantity in the *cachaça*.

The differences between experimental and simulated contents of acetic acid could be explained by the fact that the experimental volatile acidity should be, according to the Legislation, determined by titration and expressed only in terms of acetic acid. However, there are other organic and inorganic acids present in *cachaça*, and not considered in the simulation. Those acids surely contributed to increase the experimental values. Besides acetic acid, there are other volatile organic acids that are normal byproducts of the alcoholic fermentation, such as formic, butyric, and propionic acids, present in small quantities (Cardoso *et al.*, 2003; Nascimento *et al.*, 1998; Nykänen and Nykänen, 1991; Amerine *et al.*, 1972). For instance, formic acid is more volatile than acetic acid, so it will distill first, increasing the distillate acidity. On the other hand, volatile inorganic acids, such as carbonic and sulfurous acids can also be present in the wine. Carbon dioxide is produced during fermentation and sulfur dioxide can be used for sugar bleaching, so that residues of these volatile components may be present in *cachaça*, also contributing for the higher acidity values observed in the experimental runs.

In order to improve the understanding of the volatile acidity behavior during alembic distillation, two actions should be considered in future investigations. In the case of the experimental runs, a series of other acids should be monitored, since prior investigations have detected their presence in spirits (Cardoso *et al.*, 2003; Nascimento *et al.*, 1998; Nykänen and Nykänen, 1991; Amerine *et al.*, 1972). In relation to the simulation approach, those acidic compounds should be incorporated in the software

and the capacity of the equilibrium modeling to correctly predict their volatility behavior should also be evaluated.

As cited previously, the experimental distillates were separated in cuts: the head corresponded to the initial 200 mL of distillate, while the heart ended when the distillate reached about 38% v/v ethanol. The only experiment that did not present a spirit within specifications for all the compounds was W₁. In the case of W₁, the ethyl acetate content (205.8 mg/100 mL AA) was above the Legislation limit (200 mg/100 mL AA). As should be noted, W₁ had its initial ethyl acetate composition changed by a significant amount. In fact, if the concentrations of several minor compounds in the wine were close to or above the maximum values reported in the literature, the use of a simple distillation process may not be able to produce spirits within the required quality limits. In this case, the use of a batch distillation with reflux would probably be the best alternative in order to improve the separation of minor compounds in specific cuts.

The same cuts were made in the W₂ and W₃ simulations, maintaining the experimental cut times. Table 5 presents the experimental and simulated cuts. The W₂ simulation showed a *cachaça* within the Legislation limits, and concentrations close to the ones obtained experimentally, with exception of acetic acid and ethyl acetate in W₂. For acetic acid, it was observed that its concentration in the simulated spirits (2.6 mg/100 mL AA) was much lower than the experimental ones (101.6 mg/100 mL AA). This was expected, since in the profiles (Figure 7) it can be seen that the program provided a much lower content than in the experimental distillate, despite the fact that the initial acid content was modified in all experimental runs.

Table 5: Experimental and simulated cut compositions

Data	Experimental				Simulation			
	W ₂		W ₃		W ₂		W ₃	
	Head	Heart	Head	Heart	Head	Heart	Head	Heart
Time (s)	690	4876	706	4534	690	4876	706	4534
GL _{accumulated} (% v/v)	54.4	38.1	54.1	38.9	54.2	40.1	54.0	40.7
Composition (mg/100 mL AA)								
Acetaldehyde	7.1	0.5	10.0	1.3	9.4	1.6	32.4	5.6
Ethyl Acetate	150.9	21.2	154.6	21.7	518.5	50.4	219.1	21.8
Methanol	15.2	16.7	15.9	16.8	12.6	16.0	20.1	25.2
n-Propanol	66.8	54.3	73.6	59.8	70.9	51.8	103.2	76.6
Isobutanol	101.0	53.0	109.6	56.5	98.8	53.8	157.3	87.5
Isoamyl alcohol	548.9	234.1	536.4	228.4	191.1	228.6	181.2	215.2
Acetic Acid	60.9	101.6	56.6	108.6	1.8	2.6	1.7	2.4
Sum Higher Alcohols	716.7	341.4	719.6	344.7	360.7	334.3	441.7	379.4

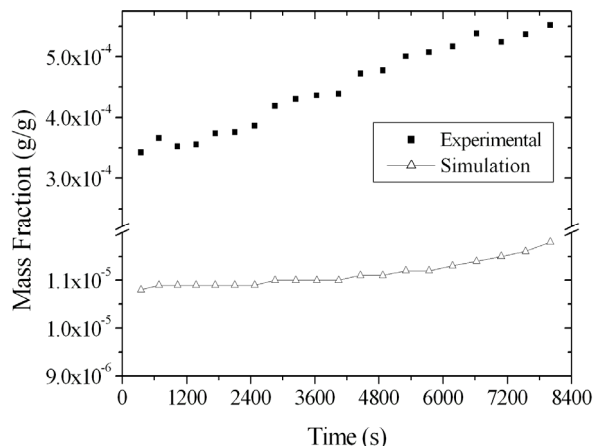


Figure 7: Distillate Volatile Acidity profiles, in acetic acid for W_2 distillation.

Unfortunately, in the case of W_3 the simulation results for methanol (25.2 mg/100 mL AA) and for the sum of higher alcohols (379.4 mg/100 mL AA) are outside the Legislation limits (20 and 360 mg/100 mL AA, respectively). In the case of the higher alcohols, only the isoamyl alcohol initial content was modified and its profiles were not well described by the simulation. However, the content of these compounds in the simulated heart cut is not too different from the experimental one. For methanol, as said before, the simulation predicted a higher concentration in the distillate than in the experimental run (see Figure 5). It should also be noted that the methanol content in W_3 is also higher than in W_2 (see Table 1).

CONCLUSION

Although the distillation path of the minor compounds is very important for spirits quality, their experimental behavior is not yet fully understood, as in the case of methanol, and their reproduction by simulation tools is also not completely satisfactory. In the present work, a series of experiments conducted with diluted and modified *cachaças* allows the measurement of heat transfer coefficients during distillation and the reliable monitoring of the minor compounds' distillation path. The suggested simulation algorithm was able to reproduce correctly the distillate amount, the wine boiling temperature, the distillate ethanol content and the concentration profiles of most minor components. The experimental and simulated results for methanol do not confirm prior data reported in the literature that indicated an unexpected increase in the methanol

concentration at the end of the distillation. The experimental profiles for isoamyl alcohol and volatile acidity were not well reproduced by the simulation. A better prediction of the vapor-liquid equilibrium for isoamyl alcohol and the inclusion of further acidic components in the wine composition are probably required in order to improve the simulation tools. Despite this drawback, the simulation provides reliable results for most compounds and can help define appropriate distillate cuts for guaranteeing spirits quality.

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NOMENCLATURE

A	heat transfer area	m^2
D	total amount of distillate	mol
D_i	total amount of component i in the distillate	mol
H_L	liquid phase enthalpy	J/mol
H_V	vapor phase enthalpy	J/mol
L	total amount of liquid in the still	mol
P	total pressure	Pa
P_{vpi}	vapor pressure of component i	P
Q	heat transferred to the boiling wine	J/s
t	time	s

T_b	wine boiling temperature	°C
T_u	heat source temperature evaluated as the temperature under the still	°C
U	heat transfer coefficient	W/m ² °C
v	vaporization rate	mol/s
x_i	liquid phase mole fraction of component i	
y_i	vapor phase mole fraction of component i	

Greek Letters

γ_i	activity coefficient of component i
ϕ_i	fugacity coefficient of component i

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